Additional data file 7. List of genes differentially expressed in pharyngeal cell-adherent pneumococci vs. culture medium control pneumococci. Numbers in bold type represent genes that met the criteria of being both statistically differentially expressed and having a ratio of  $\geq 2$  or  $\leq 0.5$  in both strains. This gene list was supplemented with genes that did not meet the fold threshold value but appeared to be co-regulated as part of an operon. NA indicates data points that were removed after analysis with Spotfinder. X indicates genes that did not meet the criteria established for microarray analysis in MeV (see methods).

		Gene	Fold increase	or decrease G54 adherent/	
Locus	Common Name	Symbol	control	control Fur	nction
SP_0014 SP_0018	Transcriptional regulator ComX1 Hypothetical protein	comX1	X 1.7	2.9 Cell	lular processes/pathogenesis/Regulatory function pothetical proteins
SP 0019	Adenylosuccinate synthetase	purA	1.0	1.7 Pur	ines, pyrimidines, nucleosides, and nucleotides known protein
SP_0020	Cytidine-deoxycytidylate deaminase family protein	•	1.3	2.5 Unk	known protein
SP_0024 SP_0025	Conserved hypothetical protein Hypothetical protein		2.6 2.7	0.9 Hyp	pothetical proteins pothetical proteins
SP_0026	Hypothetical protein		2.7	X Hvo	pothetical proteins
SP 0042	Transport atp-binding protein ComA	comA	X	2.4 Cell	Jular processes/pathogenesis/Transport and binding proteins
SP_0043	Transport protein	comB	X	2.6 Cell	lular processes/pathogenesis/Transport and binding proteins
SP_0044 SP_0045	Phosphoribosylaminoimidazole-succinocarboxamide synthase Phosphoribosylformylglycinamidine synthase II	purC purL	0.2 <b>0.2</b>	0.6 Cell	lular processes/pathogenesis lular processes/pathogenesis/Purines, pyrimidines, nucleosides,
SP_0046	Amidophosphoribosyltransferase	puiL	0.4	0.4 Cen	ines, pyrimidines, nucleosides, and nucleotides
SP 0047	Phosphoribosylformylglycinamide cyclo-ligase		0.4	0.5 Pur	ines, pyrimidines, nucleosides, and nucleotides
SP_0048	Phosphoribosylglycinamide formyltransferase		0.5	<b>0.4</b> Pur	ines, pyrimidines, nucleosides, and nucleotides
SP_0049 SP_0050	Vanz protein, putative Phosphoribosylaminoimidazolecarboxamide	purH	0.6 <b>0.5</b>	0.5 Unk	known protein ines, pyrimidines, nucleosides, and nucleotides
SP_0050	formyltransferase-imp cyclohydrolase	purn	0.5	0.4 Pur	ines, pyrimidines, nucleosides, and nucleotides
SP_0051	Phosphoribosylamineglycine ligase		0.3	0.4 Pur	ines, pyrimidines, nucleosides, and nucleotides
SP_0052	Hypothetical protein	_	0.3	<b>0.4</b> Hyp	pothetical proteins
SP_0053 SP_0054	Phosphoribosylaminoimidazole carboxylase, catalytic subunit Phosphoribosylaminoimidazole carboxylase, ATPase subunit	purE	0.4 0.4	0.5 Cell	lular processes/pathogenesis/Purines, pyrimidines, nucleosides, lular processes/pathogenesis/Purines, pyrimidines, nucleosides,
SP_0054 SP_0071	Zinc metalloprotease ZmpC	zmpC	0.6	0.4 Cell	lular processes/pathogenesis/Protein fate
SP 0083	DNA-binding response regulator	2,00	0.5	1.0 Sign	nal transduction
SP 0084	Sensor histidine kinase		0.4	0.8 Sign	nal transduction
SP_0099	Hypothetical protein		2.2	1.2 Hyp	pothetical proteins
SP_0100 SP_0101	Conserved hypothetical protein Putative transporter		2.2 1.4	1.4 Hyp	pothetical proteins nsport and binding proteins
SP_0103	Putative capsular polysaccharide biosynthesis protein		1.6	2.4 Cell	I envelope
SP_0107	LysM domain protein		1.4	2.7 Cell	I envelope
SP_0110	Bacteriocin-associated integral membrane protein		0.8	0.6 Cell	lular processes/pathogenesis
SP_0111 SP_0115	Putative amino acid ABC transporter, ATP-binding protein Hypothetical protein		1.1 0.8	0.6 Trai	nsport and binding proteins pothetical proteins
SP_0124	Hypothetical protein		0.0 X	3.3 Hyp	pothetical proteins
SP_0126 SP_0139	Hypothetical protein		X	1.9 Hyp	oothetical proteins
SP_0139	Conserved domain protein		X	0.4 Hyp	oothetical proteins
SP_0140	Udp-glucose 6-dehydrogenase, authentic frameshift	ugd	X	0.4 Cell	l envelope
SP_0142	Hypothetical protein Conserved domain protein		1.3 1.3	0.7 Hyp	pothetical proteins pothetical proteins
SP_0143 SP_0144 SP_0145	Hypothetical protein		1.8	0.6 Hyp	pothetical proteins
SP_0145	Conserved hypothetical protein		1.6	0.7 Hyp	oothetical proteins
SP_0146 SP_0147	Conserved hypothetical protein		1.3	0.7 Hyp	oothetical proteins
SP_0147	Hypothetical protein ABC transporter, substrate-binding protein		1.4 0.4	0.6 Hyp	pothetical proteins nsport and binding proteins
SP_0148 SP_0149	Lipoprotein		0.4 X	0.7 Ha	I envelope
SP_0158 SP_0159	Nrdi family protein		1.8 0.7		known protein
SP_0159	Conserved hypothetical protein		0.7	0.3 Hyp	oothetical proteins
SP_0169	Lactose phosphotransferase system repressor, degenerate		2.1	NA Disi	rupted reading frame
SP_0170	Hypothetical protein Conserved hypothetical protein		2.1 2.3	NA Hyp	pothetical proteins pothetical proteins
SP_0169 SP_0170 SP_0181 SP_0267	N5.n10-methylenetetrahydromethanopterin reductase homolog		0.3	X Unk	known protein
SP_0282 SP_0283	PTS system, mannose-specific IID component PTS system, mannose-specific IIC component PTS system, mannose-specific IIAB components		1.0	0.5 Trai	nsport and binding proteins/Signal transduction nsport and binding proteins/Signal transduction
SP_0283	PTS system, mannose-specific IIC component	manM	0.8	0.5 Trai	nsport and binding proteins/Signal transduction
SP_0284 SP_0285	Alcohol dehydrogenase, propanol-preferring	manL	0.8 <b>0.4</b>	0.3 Irai	nsport and binding proteins/Signal transduction ergy metabolism
SP_0287 SP_0288	Xanthine-uracil permease family protein		0.2	0.4 Trai	nsport and binding proteins
	Xanthine-uracil permease family protein Conserved hypothetical protein		0.2 0.4	<b>0.4</b> Hyp	nsport and binding proteins pothetical proteins
SP_0289	Dihydropteroate synthase		0.6	0.6 Bios	synthesis of cofactors, prosthetic groups, and carriers/
SP 0290	Dihydrofolate synthetase.		0.7	0.5 Pior	lular processes/patnogenesis/Energy metabolism
SP_0291	GTP cyclohydrolase I		0.7	0.5 Bios	synthesis of coractors, prosthetic groups, and carriers lular processes/pathogenesis/Energy metabolism synthesis of cofactors, prosthetic groups, and carriers synthesis of cofactors, prosthetic groups, and carriers
SP_0292 SP_0366	Bifunctional folate synthesis protein		0.6	0.5 Bios	synthesis of cofactors, prosthetic groups, and carriers
SP_0366	Oligopeptide-binding protein alia precursor (exported protein 1) Conserved hypothetical protein		1.0	0.3 Trai	synthesis of colactors, prostnetic groups, and carners nsport and binding proteins bothetical proteins
SP_0371 SP_0375	Conserved hypothetical protein		2.0 0.5	X Hyp	pothetical proteins
SP_0375 SP_0409	6-phosphogluconate dehydrogenase, decarboxylating Conserved hypothetical protein		0.6	0.6 Ene	ergy metabolism
SP 0415	Enoyl-coa hydratase	phaB	1.5	0.4 Fatt	pothetical proteins ty acid and phospholipid metabolism
SP_0416	Transcriptional regulator, MarR family	marR	1.7	X Rec	gulatory function
SP_0417	3-oxoacyl-(acyl-carrier-protein) synthase III		1.6	1.0 Fatt	ty acid and phospholipid metabolism ty acid and phospholipid metabolism ty acid and phospholipid metabolism
SP_0418 SP_0419	Acyl carrier protein Enoyl-(acyl-carrier-protein) reductase	fabK	1.8 1.4	0.9 Fatt	ty acid and phospholipid metabolism ty acid and phospholipid metabolism
SP 0420	Malonyl coa-acyl carrier protein transacylase	iaur	1.9	0.6 Fatt	tv acid and phospholipid metabolism
SP 0421	3-oxoacyl-facyl-carrier protein) reductase		2.7	0.7 Fatt	ty acid and phospholipid metabolism
SP_0422	3-oxoacyl-(acyl-carrier-protein) synthase II Acetyl-CoA carboxylase, bitoin carboxyl carrier protein	fabF	2.4	0.7 Fatt	ty acid and phospholipid metabolism
SP_0423 SP_0424	Acetyl-CoA carboxylase, bitoin carboxyl carrier protein Similar to hydroxymyristoyl-(acyl carrier protein) dehydratase.		3.6 3.6	0.9 Fatt	y acid and phospholipid metabolism ty acid and phospholipid metabolism
SP_0424 SP_0425	Acetyl-coa carboxylase, biotin carboxylase		3.6 4.2	1.0 Fatt 1.1 Fatt	ty acid and phospholipid metabolism ty acid and phospholipid metabolism
SP 0426	Acetyl-coa carboxylase, biotin carboxylase Acetyl-coa carboxylase, carboxyl transferase subunit beta	accD	5.5	1.4 Fatt	ty acid and phospholipid metabolism
SP 0427	Acetyl-coa carboxylase, carboxyl transferase subunit alpha	accA	2.5	1.6 Fatt	ty acid and phospholipid metabolism
SP_0429 SP_0430	Hypothetical protein		X X	2.8 Hyp	pothetical proteins pothetical proteins
SP_0430 SP_0445	Hypothetical protein Acetolactate synthase, large subunit, biosynthetic type	ilvB	0.8	4.1 Hyp 0.4 Δmi	no acid biosynthesis
SP 0446	Acetolactate synthase, small subunit	ilvN	0.6	0.2 Ami	ino acid biosynthesis
SP 0447	Ketol-acid reductoisomerase	ilvC	0.7	0.2 Ami	ino acid biosynthesis
SP_0448	Hypothetical protein		0.8	0.2 Hyp	pothetical proteins
SP_0449 SP_0450	Hypothetical protein Threonine dehydratase	ilvA	0.8 1.0	0.2 Hyp	pothetical proteins ino acid biosynthesis
SP_0450 SP_0461	Transcriptional regulator, putative	iIVA	1.0 2.1	U.4 AMI NA Cell	ino acid biosynthesis lular processes/pathogenesis/Regulatory function
SP 0462	Cell wall surface anchor family protein		2.6	NA Cell	I envelope/Cellular processes/pathogenesis
SP 0463	Cell wall surface anchor family protein		1.9	NA Cell	I envelope/Cellular processes/pathogenesis
SP_0464	Cell wall surface anchor family protein		2.1	NA Cell	I envelope/Cellular processes/pathogenesis
SP_0466 SP_0467	Sortase, putative Sortase, putative		2.0 2.3	NA Cell	I envelope/Cellular processes/pathogenesis/Protein fate I envelope/Cellular processes/pathogenesis/Protein fate
			2.3	NA Cell	l envelope/Cellular processes/pathogenesis/Protein fate
SP 0468	Sortase, putative		22		
SP_0468 SP_0494 SP_0496	Sortase, putative Ctp synthetase. Na/Pi cotransporter II-related protein		2.2 1.2 1.3	2.2 Pur	rines, pyrimidines, nucleosides, and nucleotides known protein

	Fold increase or decrease						
Locus	Common Name	Gene Symbol	TIGR4 adherent/ control	G54 adherent/ control	- Function		
SP_0528	Peptide pheromone BlpC	blpC	Х	0.5	Cellular processes/pathogenesis/Transcription		
SP_0529 SP_0530	Transport protein BlpB Transport ATP-binding protein ComA	blpB comA	X X	0.4	Cellular processes/pathogenesis/Transport and binding proteins		
SP_0535	Hypothetical protein  Hypothetical protein	COITIA	x		Disrupted reading frame Hypothetical proteins		
SP_0539	Bacteriocin BlpM	blpM	X	0.5	Cellular processes/pathogenesis		
SP_0540	BlpN protein	blpN	X	0.5	Cellular processes/pathogenesis		
SP_0541 SP_0543	Bacteriocin BlpO Hypothetical protein	blpO	1.1 X		Cellular processes/pathogenesis Hypothetical proteins		
SP 0544	Immunity protein BlpX	blpX	X	0.6	Cellular processes/pathogenesis		
SP_0545	Immunity protein BlpY	blpY	X		Cellular processes/pathogenesis		
SP_0607 SP_0608	Amino acid abc transporter, permease protein ABC transporter membrane-spanning permease	glnP	X 1.4		Transport and binding proteins Transport and binding proteins		
SP_0609	Amino acid ABC transporter, amino acid-binding protein	giiii	X		Cellular processes/pathogenesis/Transport and binding proteins		
SP_0610	Amino acid ABC transporter, ATP-binding protein		1.3	1.8	Transport and binding proteins		
SP_0617 SP_0626	Conserved domain protein  Branched-chain amino acid transport system II carrier protein	bmQ	1.5 1.0	3.6	Hypothetical proteins Transport and binding proteins		
SP 0685	Hypothetical protein	DITIQ	1.0 X	0.5	Hypothetical proteins		
SP_0686	Bacteriocin-associated integral membrane protein		X	0.4	Cellular processes/pathogenesis		
SP_0696	Hypothetical protein		2.1		Hypothetical proteins		
SP_0697 SP_0701	Abc transporter, atp-binding protein, authentic point mutation Orotidine 5'-phosphate decarboxylase		1.8 1.0	1NA 2.2	Transport and binding proteins Purines, pyrimidines, nucleosides, and nucleotides		
SP_0702	Orotate phosphoribosyltransferase PyrE		X	2.0	Purines, pyrimidines, nucleosides, and nucleotides		
SP 0726	Phosphomethylpyrimidine kinase		0.3	1.1	Biosynthesis of cofactors, prosthetic groups, and carriers		
SP_0730 SP_0737	Pyruvate oxidase Sodium-dependent transporter		0.3 2.4		Cellular processes/pathogenesis/Energy metabolism Cellular processes/pathogenesis/Transport and binding proteins		
SP 0738	Conserved domain protein		2.0	x	Hypothetical proteins		
SP_0753	Branched-chain amino acid ABC transporter, ATP-binding protein.		1.3	0.5	Transport and binding proteins		
SP_0766 SP_0784	Manganese co-factored superoxide dismutase.		0.3 0.1	0.5	Cellular processes/pathogenesis  Biosynthesis of cofactors, prosthetic groups, and carriers/Cellular pro		
SP_0786	Glutathione reductase ABC transporter ATP-binding protein - unknown substrate		0.1	2.9	Transport and binding proteins		
SP 0798	Dna-binding response regulator ciar	ciaR	1.2	2.9	Regulatory function/Signal transduction		
SP_0799	Sensor histidine kinase CiaH	ciaH	1.3	3.1	Regulatory function/Signal transduction		
SP_0820 SP_0845	ATP-dependent Clp protease, ATP-binding subunit ClpE Lipoprotein	clpE	0.4 <b>0.5</b>	0.9	Cellular processes/pathogenesis/Protein fate Cell envelope		
SP 0856	Branched-chain amino acid aminotransferase	ilvE	1.3	0.5	Amino acid biosynthesis		
SP_0857	Oligopeptide-binding protein, internal deletion, authentic point mutation	1	1.3		Disrupted reading frame		
SP_0858 SP_0860	Membrane protein Pyrrolidone-carboxylate peptidase		1.3 1.4	0.5	Cell envelope Protein fate		
SP_0867	ABC transporter, ATP-binding protein		0.3		Transport and binding proteins		
SP_0868	Conserved hypothetical protein		0.3	0.5	Hypothetical proteins		
SP_0869 SP_0870	YurW protein NifU family protein		0.3 0.3	0.5	Unknown protein Unknown protein		
SP_0879	Hypothetical protein		2.0		Hypothetical proteins		
SP 0999	Cytochrome c-type biogenesis protein CcdA	ccdA	0.3	0.6	Energy metabolism		
SP_1000 SP_1003	Thioredoxin family protein	-40	0.2	0.4	Energy metabolism		
SP_1003 SP_1004	Conserved hypothetical protein Hypothetical protein.	phtD	0.8 X	3.9	Hypothetical proteins Hypothetical proteins		
SP 1014	Dihydrodipicolinate synthase	dapA	1.0	0.4	Amino acid biosynthesis		
SP_1027 SP_1045	Conserved hypothetical protein		1.7	6.0	Hypothetical proteins		
SP_1045 SP_1128	Conserved hypothetical protein Phosphopyruvate hydratase	eno	0.4 0.4	0.9	Hypothetical proteins Energy metabolism		
SP 1154	Immunoglobulin A1 protease	CHO	0.5		Cellular processes/pathogenesis/Protein fate		
SP_1154 SP_1174	Conserved domain protein		X	3.3	Hypothetical proteins		
SP_1175	Conserved domain protein		1.1 0.5		Hypothetical proteins		
SP_1226 SP_1227	Histidine kinase DNA-binding response regulator		0.5	0.9	Signal transduction Signal transduction		
SP_1227 SP_1228	YfhQ protein		0.3	0.9	DNA metabolism		
SP 1229	Formatetetrahydrofolate ligase		0.3	0.9	Central intermediary metabolism		
SP_1230 SP_1231	Conserved hypothetical protein Flavoprotein		0.5 0.5	1.2	Hypothetical proteins Unknown protein		
SP_1241	Amino acid ABC transporter, amino acid-binding protein/permease pro	tein	1.6	2.7	Transport and binding proteins		
SP_1242	Amino acid ABC transporter, ATP-binding protein		0.5	2.7	Transport and binding proteins		
SP_1252 SP_1253	Hypothetical protein Hypothetical protein		X X		Hypothetical proteins Hypothetical proteins		
SP 1254	Hypothetical protein		X	1.9	Hypothetical proteins		
SP_1255 SP_1256	Putative 3-isopropylmalate dehydratase, small subunit		X	2.1	Amino acid biosynthesis		
SP_1256 SP 1257	Conserved hypothetical protein 3-isopropylmalate dehydrogenase, authentic point mutation	leuB	X	1.9	Hypothetical proteins Amino acid biosynthesis		
SP 1258	Putative 2-isopropylmalate synthase	ICUD	X		Amino acid biosynthesis		
SP_1259 SP_1260	Conserved hypothetical protein		X	1.7	Hypothetical proteins		
SP_1260	Copper homeostasis protein, CutC	cutC	1.2	1.8	Transport and binding proteins		
SP_1261 SP_1266	Conserved hypothetical protein DNA processing protein DprA, putative		1.5 X		Hypothetical proteins Cellular processes/pathogenesis		
SP_1266 SP_1267	LicC protein	licC	4.0	3.4	Cell envelope		
SP_1268	LicB	licB	4.0		Cell envelope		
SP_1269	Choline kinase. Alcohol dehydrogenase, zinc-containing		3.1 3.1	3.6	Cell envelope Cell envelope/Energy metabolism		
SP_1270 SP_1271	Putative 2-C-methyl-D-erythritol 4-phosphate cytidylyltransferase		2.1	2.9	Biosynthesis of cofactors, prosthetic groups, and carriers		
SP_1272	Repeating unit transporter.		1.6	1.9	Cell envelope		
SP_1273 SP_1274	LicD1	licD1	1.3		Cell envelope		
SP_1274 SP_1275	LicD2protein Carbamoyl-phosphate synthase, large subunit	licD2 carB	1.4 0.5		Cell envelope Purines, pyrimidines, nucleosides, and nucleotides		
SP_1276	Carbamoyl-phosphate synthase, small subunit		0.6		Purines, pyrimidines, nucleosides, and nucleotides		
SP_1277 SP_1294	Aspartate carbamoyltransferase	are D	0.5	1.1	Purines, pyrimidines, nucleosides, and nucleotides		
SP 1205	CrcB protein CrcB protein	crcB crcB	2.0 1.8		Unknown protein Unknown protein		
SP_1296	Putative chorismate mutase	CICD	1.9	1.2	Amino acid biosynthesis		
SP_1296 SP_1357 SP_1358	Putative chorismate mutase ABC transporter, permease/ATP-binding protein		0.9	0.4	Transport and binding proteins		
SP_1358	ABC transporter, ATP-binding protein. Peptide methionine sulfoxide reductase		0.7	0.3	Transport and binding proteins Cell envelope/Cellular processes/pathogenesis/Protein fate		
SP_1359 SP_1360 SP_1361	Homoserine kinase		0.7 <b>0.5</b>	0.5 0.5	Amino acid biosynthesis		
SP_1361	Homoserine dehydrogenase		0.6	0.4	Amino acid biosynthesis		
SP 1429	Peptidase, U32 family		2.1	2.5	Protein fate		
SP_1460 SP_1461	Probable amino-acid ATP transporter ATP-binding protein Yckl.  Amino acid ABC transporter, permease protein		0.6 0.6	0.6	Transport and binding proteins Transport and binding proteins		
SP_1463	Methylated-dnaprotein-cysteine s- methyltransferase		0.6	0.6	DNA metabolism		
SP_1464	Acetyltransferase, GNAT family		0.5		Unknown protein		

		Gene	Fold increase TIGR4 adherent/	or decrease G54 adherent/
Locus SD 1465	Common Name	Symbol	control 0.7	control Function
SP_1465 SP_1499	Hypothetical protein Bacterocin transport accessory protein		0.7 <b>0.2</b>	0.3 Hypothetical proteins     0.5 Cellular processes/pathogenesis/Transport and binding proteins
SP_1546 SP_1550	Hypothetical protein		0.1	0.4 Hypothetical proteins
SP_1550 SP_1551	Glutathione S-transferase family protein P-type ATPase - calcium transporter	pacL	1.1 0.9	0.3 Central intermediary metabolism     0.3 Transport and binding proteins
SP_1572	Surface located protein.	puon	1.1	2.2 Transport and binding proteins
SP_1586 SP_1587	Autoaggregation-mediating protein Oxalate-formate antiporter (OxIT-2).		0.4 0.2	Transcription     Transport and binding proteins
SP 1588	Oxidoreductase, pyridine nucleotide-disulfide, class I		0.1	0.2 Unknown protein
SP_1596 SP_1600	Is3-spn1, hypothetical protein, interruption		1.0	0.5 Disrupted reading frame
SP_1600 SP_1601	Putative membrane protein Conserved hypothetical protein		2.0 2.0	2.0 Hypothetical proteins 2.0 Hypothetical proteins
SP_1602	Required for expression of the phosphonate utilization	phnA	2.5	2.1 Transport and binding proteins
SP_1648	phenotype in E. coli Manganese ABC transporter, ATP-binding protein	psaB	2.8	2 9 Transport and hinding proteins
SP_1649	Manganese ABC transporter, permease protein,	psab	2.6	2.8 Transport and binding proteins 2.8 Transport and binding proteins
	putative, authentic frameshift			
SP_1650 SP_1651	Manganese abc transporter, manganese-binding adhesion liprotein Thiol peroxidase		2.0 0.3	2.5 Cellular processes/pathogenesis/Transport and binding proteins 0.3 Cellular processes/pathogenesis
SP_1653	ABC transporter, ATP-binding protein.		0.9	0.5 Transport and binding proteins
SP_1654 SP_1695	Conserved hypothetical protein		X	0.3 Hypothetical proteins
SP 1696	Xylan esterase 1 Hypothetical protein		1.1	3.3 Energy metabolism 2.5 Hypothetical proteins
SP_1708 SP_1714	Hypothetical protein		2.2	0.8 Hypothetical proteins
SP_1714 SP_1715	transcriptional regulator, GntR family ABC transporter, ATP-binding protein		1.4 1.1	Regulatory function     Transport and binding proteins
SP 1716	Conserved hypothetical protein		X	2.7 Hypothetical proteins
SP_1717 SP_1754	ABC transporter, ATP-binding protein		X	3.0 Transport and binding proteins
SP_1754 SP_1758	Conserved hypothetical protein Glycosyl transferase, group 1		1.8 2.2	0.5 Hypothetical proteins  NA Cell envelope
SP 1761	Hypothetical protein		1.9	NA Hypothetical proteins
SP_1762 SP_1770	Hypothetical protein		2.1	NA Hypothetical proteins
SP_1770 SP_1771	Glycosyl transferase, family 8 Glycosyl transferase, family 2-glycosyl transferase family 8		2.2 1.8	NA Cell envelope NA Cell envelope
SP_1774 SP_1775	Transcriptional regulator, putative		0.5	0.5 Regulatory function
SP_1775	Hypothetical protein	44	0.2	0.4 Hypothetical proteins
SP_1776 SP_1778	Thioredoxin reductase Water channel protein.	trxA	<b>0.2</b> 0.5	O.3 Energy metabolism     Transport and binding proteins
SP_1811 SP_1813	Tryptophan synthase, alpha subunit		1.9	0.7 Amino acid biosynthesis
SP_1813	N-(5'-phosphoribosyl)-anthranilate isomerase		2.0	2.0 Amino acid biosynthesis
SP_1814 SP_1845	Indole-3-glycerol phosphate synthase Exodeoxyribonuclease		1.8 0.4	Amino acid biosynthesis     DNA metabolism
SP_1848	Xanthine permease		0.3	X Transport and binding proteins
SP_1853 SP_1855	Galactokinase Dehydrogenase	galK	1.1 6.0	Senergy metabolism     Benergy metabolism
SP 1856	Transcriptional regulator, MerR family		7.6	1.1 Regulatory function
SP_1856 SP_1857	Cation efflux system protein		14.4	X Transport and binding proteins
SP_1860 SP_1861	Choline transporter ABC transporter ATP-binding protein - choline transporter	proWX proV	0.2 0.2	Transport and binding proteins     Transport and binding proteins
SP_1861 SP_1862	Hypothetical protein	prov	0.2	0.4 Hypothetical proteins
SP_1872	Ferric anguibactin-binding protein precusor fatb of V. anguillarum		X	0.3 Transport and binding proteins
SP_1887 SP_1888	Oligopeptide transport ATP-binding protein AmiF Oligopeptide ABC transporter, ATP-binding protein AmiE	amiF amiE	1.0 1.0	Transport and binding proteins     Transport and binding proteins
SP_1888 SP_1889	Oligopeptide abc transporter, permease protein AmiD	amiD	1.0	0.5 Cellular processes/pathogenesis/Transport and binding proteins
SP_1890	Oligopeptide transport system permease protein AmiC	amiC	1.0	0.5 Transport and binding proteins
SP_1891 SP_1893	Oligopeptide-binding protein AmiA precursor. Hypothetical protein	amiA	0.8 1.2	0.5 Cellular processes/pathogenesis/Transport and binding proteins 1.4 Hypothetical proteins
SP_1906	Chaperonin, 60 kDa		0.3	1.1 Protein fate
SP_1907	Chaperonin, 10 kDa Single-strand binding protein (ssb) (helix-destabilizing protein).		0.4 X	1.2 Protein fate     3.5 Cellular processes/pathogenesis/DNA metabolism
SP_1908 SP_1922	Conserved hypothetical protein		2.3	1.0 Hypothetical proteins
SP 1923	Pneumolysin	ply	3.0	0.5 Cellular processes/pathogenesis
SP_1924 SP_1925 SP_1926	Hypothetical protein Hypothetical protein		2.6 2.3	0.4 Hypothetical proteins     0.4 Hypothetical proteins
SP 1926	Hypothetical protein		1.7	0.4 Hypothetical proteins
SP 1945	Hypothetical protein		X	4.0 Hypothetical proteins
SP_1954 SP_1986	Serine protease, subtilase family, authentic frameshift Hypothetical protein		0.5 1.1	NA Protein fate 3.6 Hypothetical proteins
SP 1987	ABC transporter, ATP-binding protein		0.9	3.8 Transport and binding proteins
SP_1988 SP_2026 SP_2044	Bacteriocin-associated integral membrane protein		0.9	4.0 Cellular processes/pathogenesis
SP_2026 SP_2044	Alcohol-acetaldehyde dehydrogenase Acetate kinase		X 0.6	0.4 Energy metabolism     0.5 Energy metabolism
SP_2053	Competence protein		X	3.3 Cellular processes/pathogenesis
SP 2054	Hypothetical protein		0.3	1.3 Hypothetical proteins
SP_2055 SP_2108	Alcohol dehydrogenase Maltose-maltodextrin-binding protein precursor		<b>0.4</b> 0.7	O.4 Energy metabolism     O.4 Transport and binding proteins
SP 2109	Maltodextrin transport system permease protein MalC	malC	0.9	0.6 Transport and binding proteins
SP_2110	Maltodextrin ABC transporter, permease protein	malD	2.5	0.6 Transport and binding proteins
SP_2125 SP_2136	Conserved hypothetical protein PcpA	рсрА	1.3 <b>0.5</b>	0.4 Hypothetical proteins     2.4 Cell envelope/Cellular processes/pathogenesis
SP_2169	Zinc ABC transporter, zinc-binding lipoprotein		1.3	1.8 Cellular processes/pathogenesis/Transport and binding proteins
SP_2170	Zinc ABC transporter, permease protein	adcB	1.5	2.8 Cellular processes/pathogenesis/Transport and binding proteins
SP_2171 SP_2172	Zinc abc transporter, atp-binding protein Adc operon repressor AdcR	adcC adcR	1.4 1.6	2.6 Cellular processes/pathogenesis/Transport and binding proteins     2.4 Cellular processes/pathogenesis/Regulatory function
SP_2172 SP_2173 SP_2174	Extramembranal protein		1.5	2.9 Cell envelope
SP_2174	D-alanyl carrier protein	dItC	1.7	2.7 Cell envelope 2.4 Cell envelope/Transport and binding proteins
SP_2175 SP_2176	Integral membrane protein D-alanine-activating enzyme	dltB dltA	1.3 1.4	2.4 Cell envelope/Transport and binding proteins     2.5 Cell envelope
SP 2177	Hypothetical protein		1.2	2.3 Hypothetical proteins
SP 2187	Conserved domain protein		0.3	0.9 Hypothetical proteins
SP_2188 SP_2189	Chaperonin, 33 kDa TIM-barrel protein, putative, NifR3 family		0.5 0.5	1.3 Protein fate     1.1 Unknown protein
SP_2197 SP_2198	ABC transporter, substrate-binding protein, putative		0.4	0.4 Transport and binding proteins
SP_2198 SP_2199	ABC transporter, permease protein		0.5	X Transport and binding proteins
SP_2199 SP_2235	Conserved hypothetical protein Response regulator come.	comE	0.4 X	0.5 Hypothetical proteins     2.8 Cellular processes/pathogenesis/Regulatory function/Signal transductions
	Putative sensor histidine kinase comd	comD	2.3	2.8 Cellular processes/pathogenesis/Regulatory function/Signal transductions of Callular processes/pathogenesis/Regulatory function/Signal transductions
SP 2230				
SP_2236 SP_2237 SP_2239	Competence stimulating peptide precursor (CSP) HtrA protein	comC htrA	1.3 1.2	3.1 Cellular processes/pathogenesis     4.6 Protein fate

<sup>\* &</sup>quot;NA" indicates data points that were removed after analysis with spotfinder. "X" indicates genes that do not meet the criteria established for microarray analysis in MeV (see experimental procedures)